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TITLE: Identification of Novel Genes Affected by Gamma

Irradiation Using a Gene-Trapped Library of Human Mammary

Epithelial Cells

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13. ABSTRACT (Maximum 200 Words)

We propose that the expression of several unknown genes is affected by gamma radiation. Abnormal expression of these genes may be one of the steps in breast carcinogenesis induced by radiation. We plan to establish an assay that will allow us to screen for breast cells that contains a single mutation by gene trapping. We will be able to detect changes in the expression of a gene upon treatment with different doses of radiation. These radiation-responsive genes will be identified through the rapid amplification of cDNA ends procedure and sequenced. Cells that are affected by radiation will be isolated and analyzed to see if the changes can lead to transformation of the normal breast epithelial cell into a neoplastic cell. This assay may prove to be a powerful tool in the identification of novel genes that are affected by gamma irradiation in ethe early stages of breast cancer progression.

The 3'RACE protocol has recently been completed and thirty one genes potential genes were sequenced. Of these, six candidate genes were found. The include: human creatine kinase gene, human androgen receptor, human DORA reverse strand protein 1 (DREV1), human eukaryotic translation elongation factor 1 beta 2 (EEF1B2), human ribosomal protein L27, and human DNA clone RP11-290F20 on chromosome 20. These genes will be further analyzed for their transformation properties of human mammary epithelial cells as discussed in the statement of work.

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Table of Contents

Cover1
SF 2982
Table of Contents3
Introduction4
Body4
Key Research Accomplishments7
Reportable Outcomes7
Conclusions
References
Appendices8

INTRODUCTION

We propose that the expression of several unknown genes is affected by gamma irradiation. The subject and purpose of our research is that the abnormal expression of these genes may be one of the early steps in breast carcinogenesis induced by radiation. We plan to establish an assay that will allow us to screen for breast cells that contain a single mutation by gene trapping. We will be able to detect changes in the expression of a gene upon treatment with different doses of radiation. These radiation-responsive genes will be identified and analyzed to see if the changes can lead to transformation of the normal breast epithelial cell into a neoplastic cell. This assay may prove to be a powerful tool in the identification of novel genes that are affected by gamma irradiation in the early stages of breast cancer progression.

BODY

RESEARCH TRAINING

Ongoing training is very important throughout my predoctoral period. My department, Environmental and Radiological Health Sciences, places an important focus on training. Weekly, I attend Advanced Radiation Biology journal meetings where faculty and students interact and discuss current and relevant papers in breast cancer research and radiation effects. Each attendee presents one journal article every semester and leads the discussion. Weekly there is a Cell and Molecular Biology seminar where invited visiting speakers give a 50 minute presentation and discussion about relevant topics such as breast cancer research, cell signaling, and many more. There are also two departmental seminars I attend weekly where visiting speakers, as well as graduate students present their research. It is very important for my training that I keep current with the latest research techniques and discoveries by attending these meetings. My mentor, Dr. Robert Ullrich, is currently the Oncology Chair of the Veterinary Teaching Hospital here on campus, so I am also exposed to more clinical cancer research seminars and meetings that I attend there as well.

In October 2003 I attended the American Association for Cancer Research special conference on the Advances in Breast Cancer Research meeting in Huntington Beach, California. In February 2004 I attend the American Association for Cancer Research special conference on Radiation Biology and Cancer meeting in Dana Point, California. These meetings contributed a great deal to my overall predoctoral training by exposing me to breast cancer research scientists from all over the world. I was able to attend numerous oral and poster presentations and learn about the latest advances being made in breast cancer and radiation research.

RESEARCH PURPOSE & GOALS

We plan to identify novel genes affected by gamma irradiation and to characterize their function using a gene-trapped library of human mammary epithelial cells. We hypothesize that the mutation of these novel genes or its abnormal expression is one of the causes of early breast carcinogenesis. Mounting evidence suggests that gene products may function differently depending on cell type, developmental stage, or species. Thus, to identify novel gene(s) critical for the initiation of breast cancer, we need to study the irradiation effects of "loss of function" of a gene product in human breast epithelial cells.

The issue of how low dose gamma radiation may lead to breast cancer will be addressed by studying the genes affected by low dose gamma irradiation. We will focus on the trapped genes whose expression are immediately changed by a single dose of gamma irradiation, determine if this is a dose-dependent effect and further analyze whether this effect can lead to transformation of the breast cells.

The following are specific aims as outlined in the approved statement of work:

Specific Aim 1: To establish a high throughput assay for detection of variation in gene expression in human mammary epithelial cells using gene-trapped MCF10A clones. Specific Aim 2: To determine the effect of gamma irradiation on expression of reporter protein GFP (green fluorescent protein).

Specific Aim 3: To characterize the effect of gamma irradiation on transformation of human mammary epithelial cells.

Specific Aim 4: To identify the trapped genes affected by gamma irradiation.

RESEARCH PROGRESS

Currently, specific aim 1, specific aim 2, and specific aim 4 are completed. Specific Aim 3 is still in progress. At this point no transformation has been observed yet. Attached in the appendices are color representations of the completed construction of the gene-trapped MCF10A clonal library as seen under a fluorescent microscope. This is included in one of my PowerPoint presentations. It is clearly observed in the pictorials, that the bright green fluorescence luminating from the cells is due to the retrovirus pRET being incorporated into the genome.

A total of 192 gene-trapped clones were analyzed by the construction of a single cell assay in 96-well plates. This was done to obtain single cell clones, hopefully each representing a different trapped gene. One 96-well plate contained the pooled gene-trapped MCF10A cells sorted by flow cytometry into a GFP positive pool and the other 96-well plate contained the pooled gene-trapped MCF10A cells sorted by flow cytometry into a GFP negative pool. Graphical representations of the flow cytometry data are included in one of my power point presentations. Replica plating was then done from both of the original single cell assay plates for the following GFP expression levels to be measured at: basal, control, master, store at -80°C, 0.5 Gy, and 2.0 Gy gamma irradiated. GFP measurements were made with a microplate reader by the way of a sandwich ELISA assay. The sandwich ELISA assay was accomplished by first expanding the 96-well plates with the single cell clones into 24-well plates. These 24-well plates were then expanded further to allow for 2 wells for each single cell assay clone. This was done so that one well could be further expanded and frozen for later use and the other well would be utilized to collect the cell lysate from for the ELISA assay. All of the 24-well plates were then irradiated with 2.0 Gy from a ¹³⁷Cs source. The following antibodies were used for the sandwich ELISA assay: anti-GFP (Mouse) was the primary antibody and peroxidase IgG mouse (Rabbit) was the secondary antibody.

Graphical representations of the gene expression of GFP after 2.0 Gy gamma radiation dose from a ¹³⁷Cs source is attached as well. Here, clones that were up- or down-regulated at least 2-fold from basal readings were expanded for further analysis. The basal GFP readings of the gene-trapped clones are included in the appendices for comparison to the 2.0 Gy GFP readings. Out of the 192 clones analyzed, 92 were up- or down-regulated at least 2-fold in comparison to basal GFP readings. These clones were expanded in culture and RNA was collected for gene analysis.

There was a slight change in the order of the approved statement of work next. Specific aim 4 was undertaken prior to the characterization of the effect of gamma irradiation on the transformation of the MCF10A cells. This was felt to be an important substitution due to the fact

that the gene that had been trapped should be identified before transformation assays were undertaken. Transformation assays are very tedious and time consuming. If, for example, the gene trapped was an artifact, then the process of analyzing for transformation could be skipped.

Specific aim 4 dealt with the characterization of the trapped genes that were causing either an up- or down-regulation upon treatment with 2.0 Gy. To analyze this, the gene-trapped clones were expanded and RNA was collected by using Qiagen's RNeasy kit. The protocol for this procedure is attached in the appendices. The RNA was then reverse transcribed into cDNA and amplified by the use of the Advantage-GC cDNA polymerase kit from BD Biosciences and the 3'RACE protocol from Invitrogen. Both protocols are included in the appendices. Gene specific primers for the neomycin marker found on our pRET retrovirus and against the polyA tail of the endogenous gene were designed. After each step, reverse transcription, first strand cDNA synthesis, and second strand cDNA synthesis, agarose gels were run to verify that the gene products were of the correct size.

When a gene product was of the correct size they were PCR purified by Qiagen's PCR purification kit and transformed into One Shot competent E. coli cells via a TOPO Cloning kit from Invitrogen. The transformed clones were then added to LB media and grown overnight. Clones were growth had occurred were then subjected to Qiagen's mini prep kit to harvest the DNA. Protocols for both of these procedures can be found in the appendices. The mini prep clones are then subjected to PCR with M13 primers and run on 1.5% agarose gels. Gel electrophoresis images are provided in the appendices in one of my PowerPoint presentations to illustrate which clones were selected to be sequenced.

Sequencing of the positive mini prep clones was completed at Davis Sequencing which is located at the University of California at Davis. A total of 31 clones were sent off for sequencing and six yielded positive results. The six genes were determined by plugging the sequences of my clones into BLAST and searching for homologous genes. The other clones were determined to be artifacts of the cloning vector. The genes that were trapped were: human creatine kinase gene, human DORA reverse strand protein 1 (DREV1), human eukaryotic translation elongation factor 1 beta 2 (EEF1B2), human ribosomal protein L27, and human DNA clone RP11-290F20 on chromosome 20.

The genes identified through sequencing analysis were expanded for RNA collection and analyzed by real-time PCR. This experiment was performed in order to analyze gene expression of the genes identified through trapping in both the gene-trapped clones and in the parental MCF10A cell line with and without ionizing radiation treatment. We felt that it was important to analyze the identified gene expression levels following IR treatment in the parental cell line to verify that in fact we were in deed seeing a radiation response. These expression levels could also then be compared to the breast cancer cell line, MCF7. In the appendices I have included my real-time PCR protocol and the sequences for the primers and probes that were utilized. Real-time PCR experiments were conducted on an Applied Biosystems 7000 Sequence Detection System with the TaqMan Gold RT-PCR Kit. Also, graphical representations of the relative gene expression of the genes of interest in various time course experiments following ionizing radiation (IR) and after varying doses of IR are included. The time course experiments were conducted at 2, 4, 8, 12, 24, and 30 hours post ionizing radiation treatment with a dose of 2.0 Gy. I am currently in the process of completing time course studies for 0.5 Gy, 1.0 Gy, and 4.0 Gy which will not be done in time to submit for this annual report. All five genes illustrated a radiation response and their relative gene expression and n-fold difference in comparison to the

parental, MCF10A cell line, were analyzed. The experimental results from the above mentioned items are all included in the appendices.

KEY RESEARCH ACCOMPLISHMENTS

- Five radiation response genes were found to be homologous to known genes through a BLAST search. These genes include: human creatine kinase gene, human DORA reverse strand protein 1 (DREV1), human eukaryotic translation elongation factor 1 beta 2, human androgen receptor, human ribosomal protein L27, and human DNA clone RP11-290F20 on chromosome 20.
- Genes of interest were found to response to a 2.0 Gy dose of ionizing radiation and time course experiments were completed to find when peak expression levels following IR occurred. This was analyzed by real-time PCR.
- Cell cycle analysis was done to verify that there was not a cell cycle delay or block causing some of the large changes in expression of some of the genes that was seen.
- Real-time PCR analysis was performed to analyze the expression of our genes of interest at 0.5 Gy, 1.0 Gy, 2.0 Gy, and 4.0 Gy doses of IR.
- One of the genes of interest, DREV1, has a small gene called DORA located in intron 4 on the complement strand. Real-time PCR analysis has been completed to investigate if its gene expression is also affected by the IR doses.

REPORTABLE OUTCOMES

- The development of a gene-trapped cell library of MCF10A cells was accomplished with the retrovirus pRET.
- On March 18, 2004 I gave a 20-minute oral presentation on my research to the faculty and students of the Department of Radiological and Environmental Health Sciences. The PowerPoint slides from my presentation are given in the appendices.
- I was invited to give a poster presentation at Colorado State University for the Cell and Molecular Biology Interdisciplinary Graduate Program Graduate Student and Post Doc Poster Competition on February 27, 2004. My poster abstract is found in the appendices.
- I was invited to give a poster presentation at the American Association for Cancer Research Special Conference: Radiation Biology and Cancer. It was held from February 18th through February 22nd in Dana Point, California. My abstract can be found in the appendices.
- I was invited to give a poster presentation at the American Association for Cancer Research Special Conference: Advances in Breast Cancer Research. It was held in October 2003 in Huntington Beach, California. My abstract can be found in the appendices.
- On October 23, 2003 I gave a 20-minute oral presentation on my research to the faculty and students of the Department of Radiological and Environmental Health Sciences. The department has doctoral students give oral presentations every semester on how their research is progressing and any new findings. The PowerPoint slides from my presentation are given in the appendices.

APPENDICES

QIAGEN RNeasy Protocol Website:

http://www1.qiagen.com/literature/handbooks/PDF/RNAStabilizationAndPurification/FromAnimalAndPlantTissuesBacteriaYeastAndFungi/RNY_Mini/1016272HBRNY_062001WW.pdf

BD Biosciences Advantage GC cDNA PCR kit manual

http://www.bdbiosciences.com/clontech/techinfo/manuals/PDF/PT1580-1.pdf

Invitrogen 3'RACE system for amplification of cDNA ends manual

https://catalog.invitrogen.com/index.cfm?fuseaction=viewCatalog.viewProductDetails&sku=&productDetails&sku

Invitrogen TOPO Cloning Kit manual

http://www.invitrogen.com/content/sfs/brochures/710 021849%20 B TOPOCloning bro.pdf

QIAGEN PCR purification kit manual

http://www1.qiagen.com/literature/handbooks/PDF/DNACleanupAndConcentration/QQ_Spin/102142 2 HBQQSpin 072002WW.pdf

Applied Biosystems TaqMan Gold RT-PCR Kit

http://home.appliedbiosystems.com/ search product literature for manual

Reverse Transcription conditions:

25°C, 10 minutes

48°C, 30 minutes

95°C, 5 minutes

Real-Time PCR conditions

50°C, 2 minutes, 1 cycle

95°C, 10 minutes, 1 cycle

95°C, 15 seconds; 60°C, 1 minute, 40 cycles

Real-Time PCR primers (concentrations in reaction at 200 nM)

Igsf6F1

ACCTTCTCCGCAACCGG

Igsf6F2

TACCTTCTCCGCAACCGG

Igsf6F3

GTACCTTCTCCGCAACCGG

Igsf6R1

GCACCGTAGCGAAACCACA

AndrogenF1

CCCTGGCGGCATGGT

AndrogenF2

ACCCTGGCGGCATGGT

AndrogenF3

TACCCTGGCGGCATGGT

AndrogenR1

CCCATTTCGCTTTTGACACA

AndrogenR2

CCCATTTCGCTTTTGACACAA

AndrogenR3

GCCCATTTCGCTTTTGACA

DORAF1

GAGGCAGGGTCATCCTTGC

DORAF2

GAGCCAACTAGAGGCAGGGTC

DORAF3

GCCAACTAGAGGCAGGGTCA

DORAR1

CCCACTTGCCACCTACGTTT

DORAR2

TCCCACTTGCCACCTACGTT

DORAR3

CTCCCACTTGCCACCTACGT

CKF1

TGCTACCATGGGCACCAGT

CKF2

TTGCTACCATGGGCACCAGT

CKF3

TTGCTACCATGGGCACCAG

CKR1

GCACACACTTTCTGCCGGT

CKR2

GCACACACTTTCTGCCGGTT

CKR3

GGCACTCGGCCATGCA

EEF1B2F1

CACAATTTGCGCGCTCTCT

EEF1B2F2

CCACAATTTGCGCGCTCT

EEF1B2F3

CCACAATTTGCGCGCTC

EEF1B2R1

ACCCATGGTGTCGGCTGTA

EEF1B2R2

ACCCATGGTGTCGGCTGT

EEF1B2R3

AACCCATGGTGTCGGCTGTA

L27F1

GCCCCTACAGCCATGCTCT

L27F2

ATCGCCCCTACAGCCATG

L27F3

TCAGATCGCCCTACAGCC

L27R1

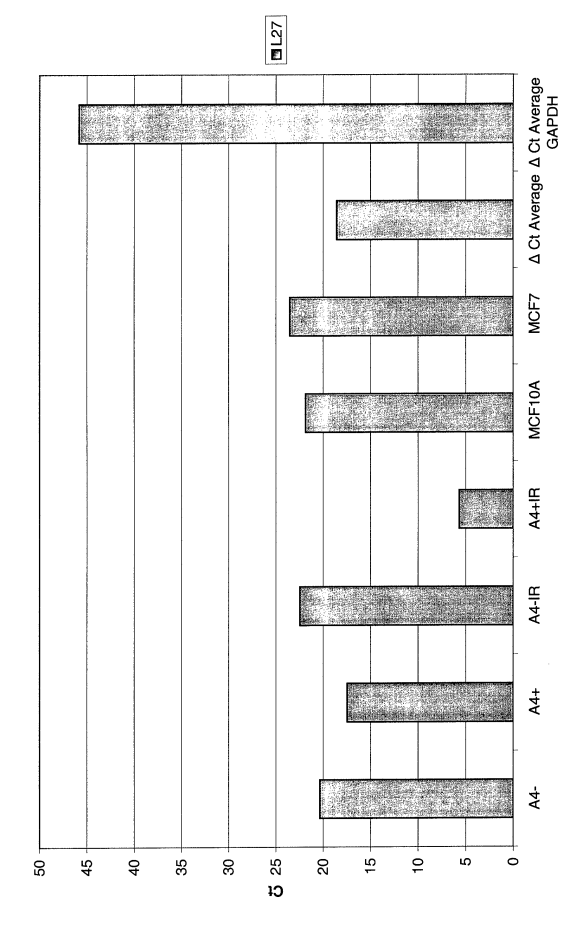
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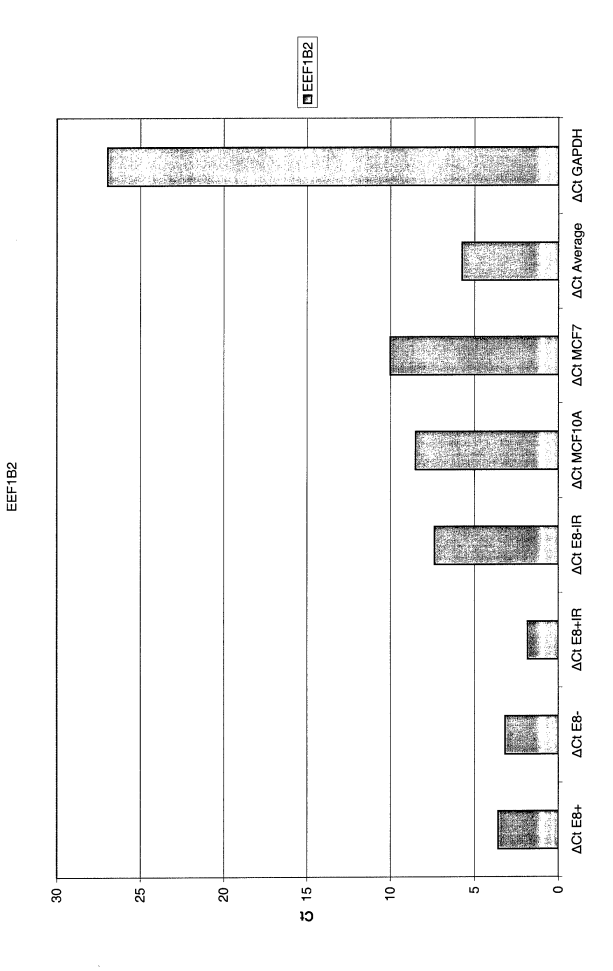
L27R2

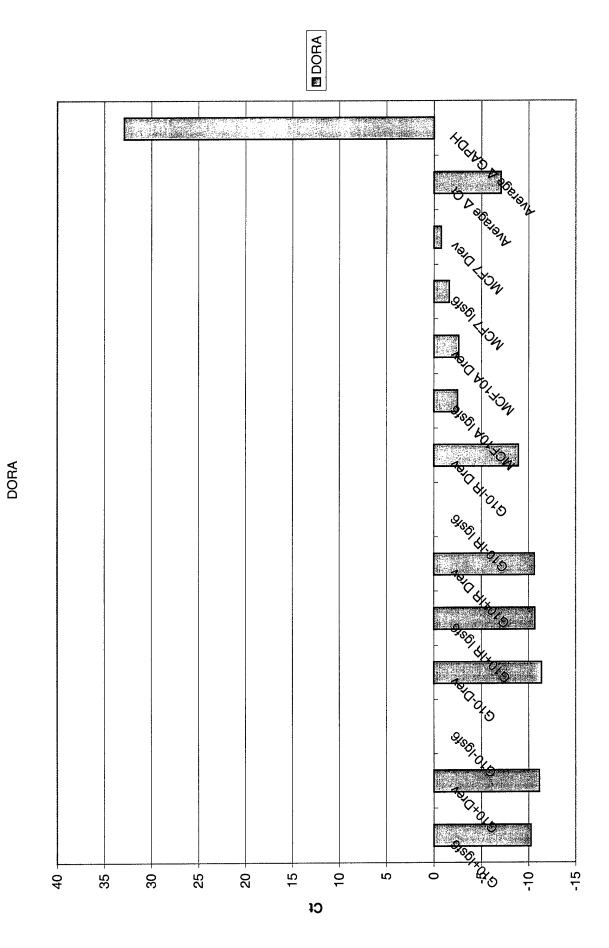
CCCATGGCAGCTGTCACTT

L27R3 TCTTGGCGATCTTCTTGC

Real-Time PCR Probes (concentration in reaction at 100nM)
Igsf6
6FAM-TGCCCTTCTGAGCAACCAACATGC-TAMRA
Androgen
6FAM-AGCAGAGTGCCCTATCCCAGTCCCA-TAMRA
DORA
6FAM-CTTGTCCTCCCCTTTCATCCCTATGTGG-TAMRA
CK
6FAM-TCCTGACCACCGGGTACCTGCTG-TAMRA
EEF1B2
6FAM-TCTGCTGCTCCCCAGCTCTCGG-TAMRA
L27
6FAM-TGGCTGGAATTGACCGCTACCCC-TAMRA







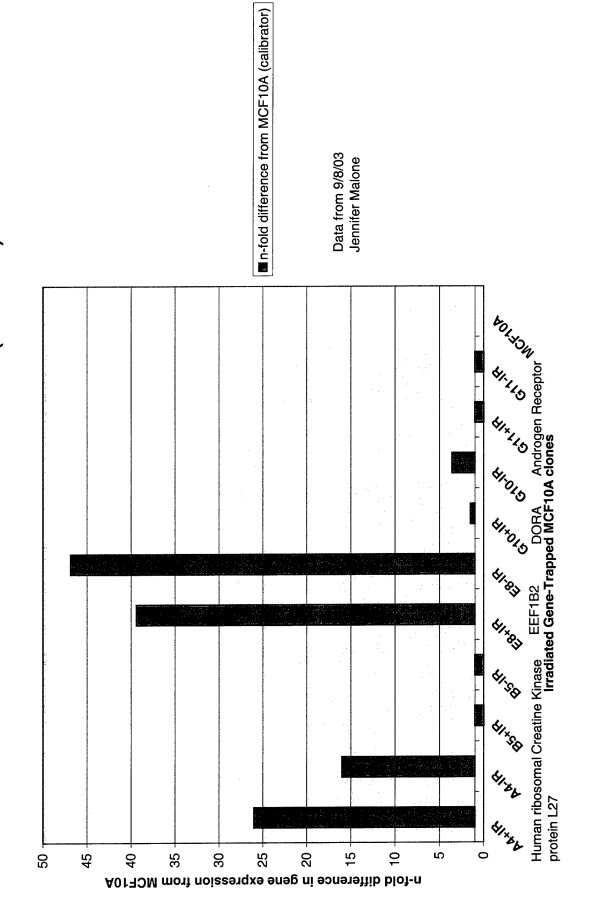
Human Creatine Kinase ACHES IR ACTINGFIOA ACTINGF7 Average A GAPDH GAPDH Ŋ -10 20 35 30 52 15 9 ٠Ş 0 ιO

Human Creatine Kinase

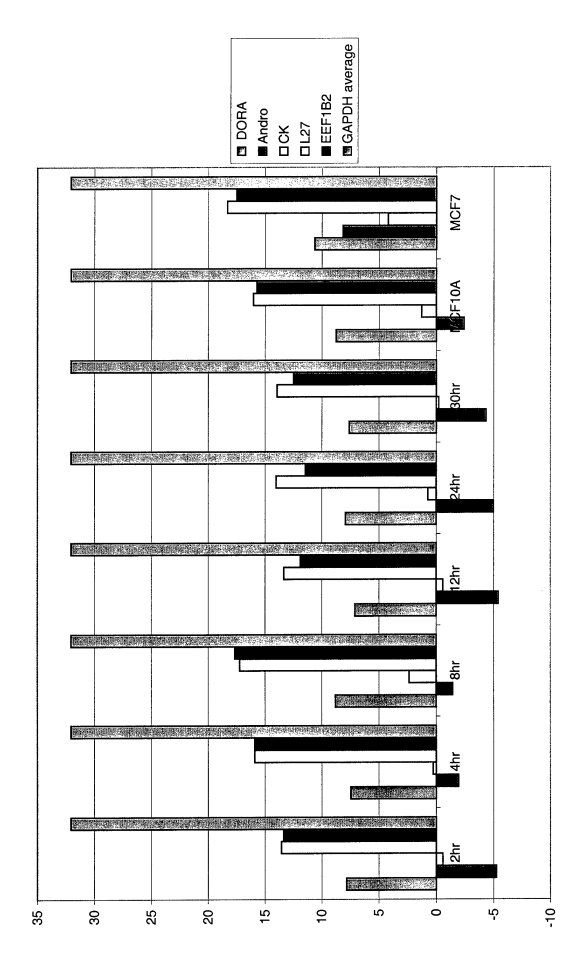
MCF10A 2hr post 2.0Gy IR

MCF10A 4hr post 2.0 Gy IR

n-fold difference from MCF10A (calibrator)



Relative Gene Expression Levels



MCF10A Post IR

Hours post IR

Data from 9/8/03 Jennifer Malone □ Creatine Kinase ■ Androgen ■ EEF1B2 ■ DORA **1**27 6 Ņ ဖ 2 က ٥ ω 4 Ţ N-fold differences

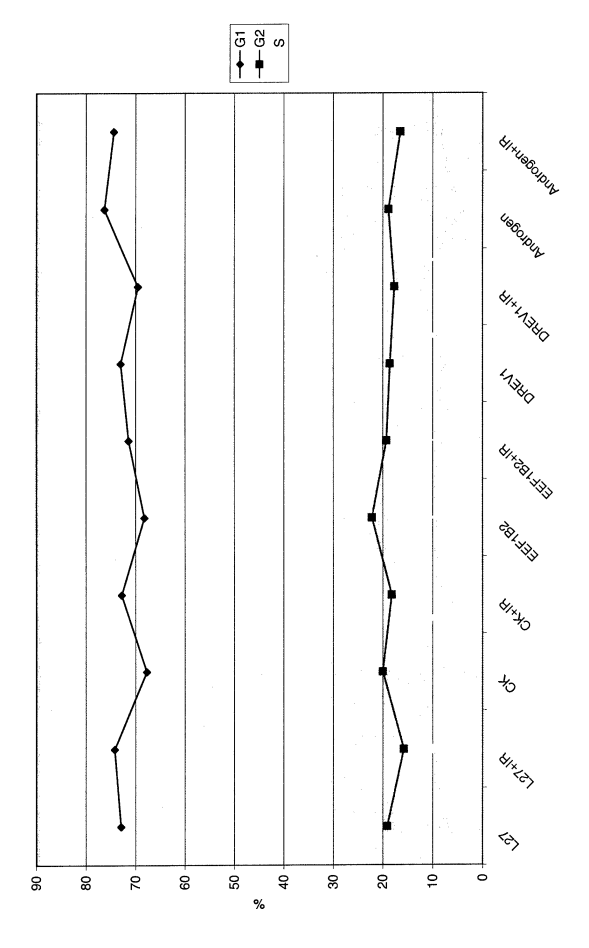
Gene Expression post IR

hours post IR

■G1 **■**G2 □S **30hr** 24hr 12hr 8hr 4hr 2hr MCF10A 20 -30 -10 -40 -0 70 -- 09 50 80 96 %

Cell Cycle Analysis post 2.0 Gy IR

Cell Cycle Analysis of Potential Radiation Response Genes

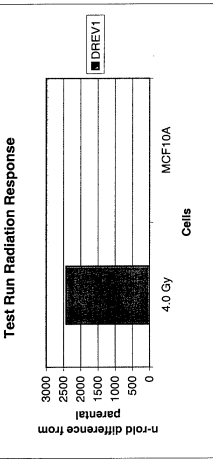


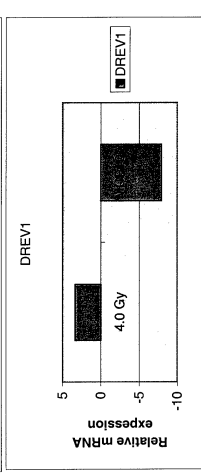
average	35.0974	31.00387	27.89373	24.4558	24,1986
10			27.8083		
	34.223	30.4173	27.8527	24.1228	23.8954
GAPDH	34.9355	31.7649	28.0202	24.781	23,3839
	80 pg	400 pg	2 ng	10 ng	50 ng
ACF10A	35.3012	37.5656		36.4334	
_			23.1694		
	DREV1	DREV1	DREV1	average	

delta Ct GAPDH 28.52988

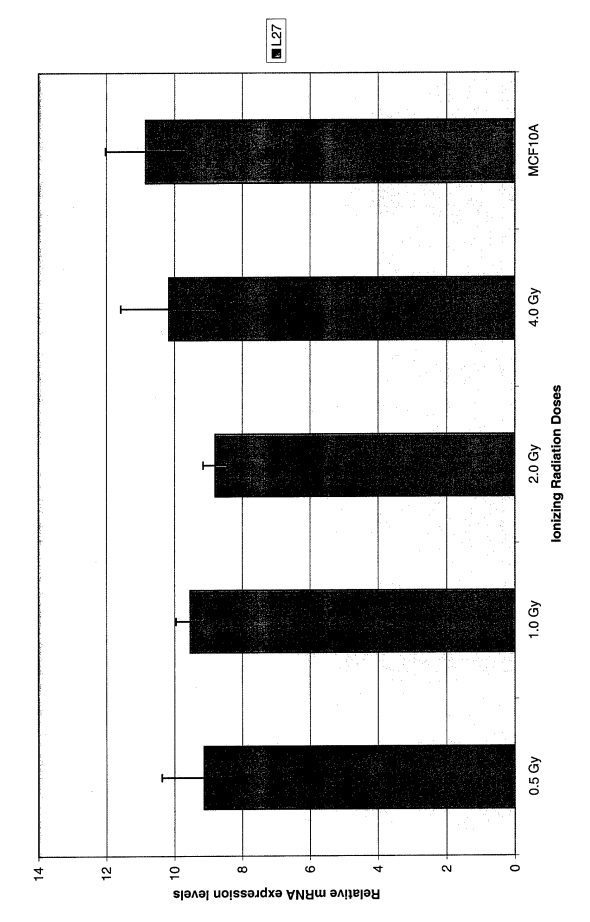
3.335747 -7.90352 deltadeltaCt 4.0 Gy 3 MCF10A

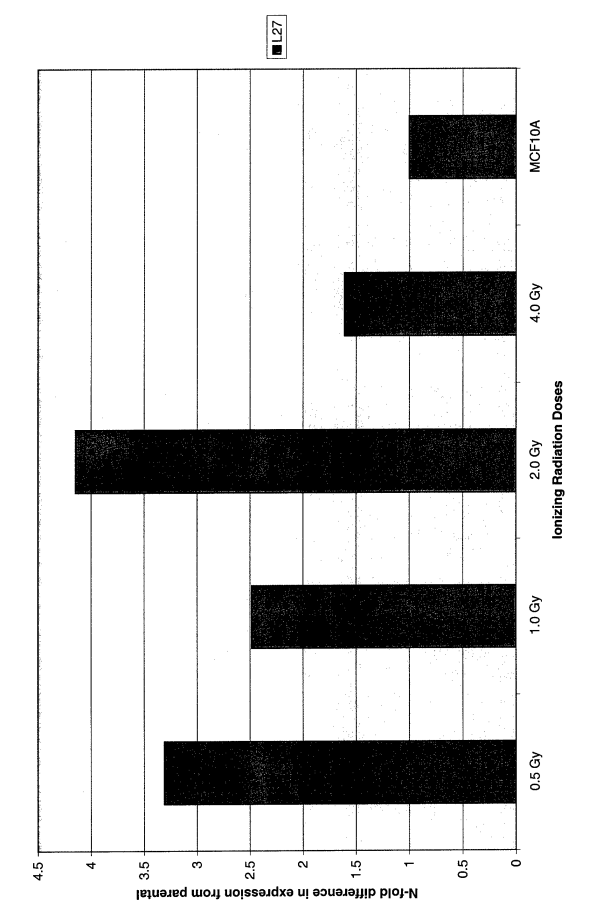
n-fold differences 4.0 Gy 2417.444 MCF10A 1



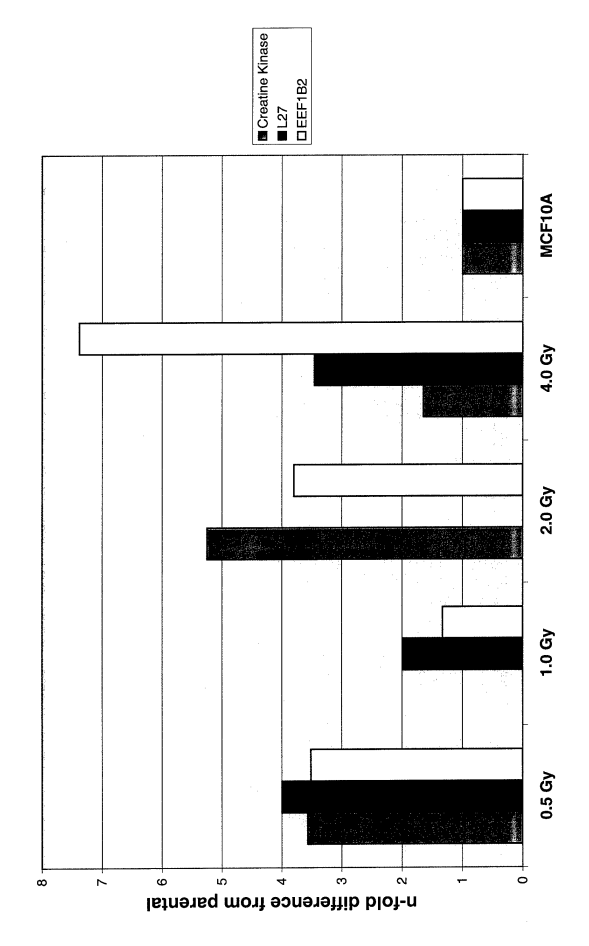


DORA

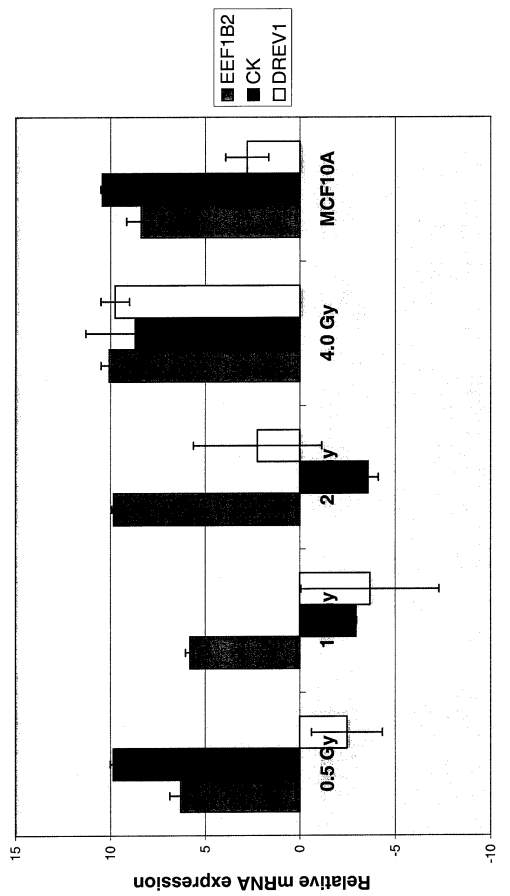




N-Fold Differences from MCF10A

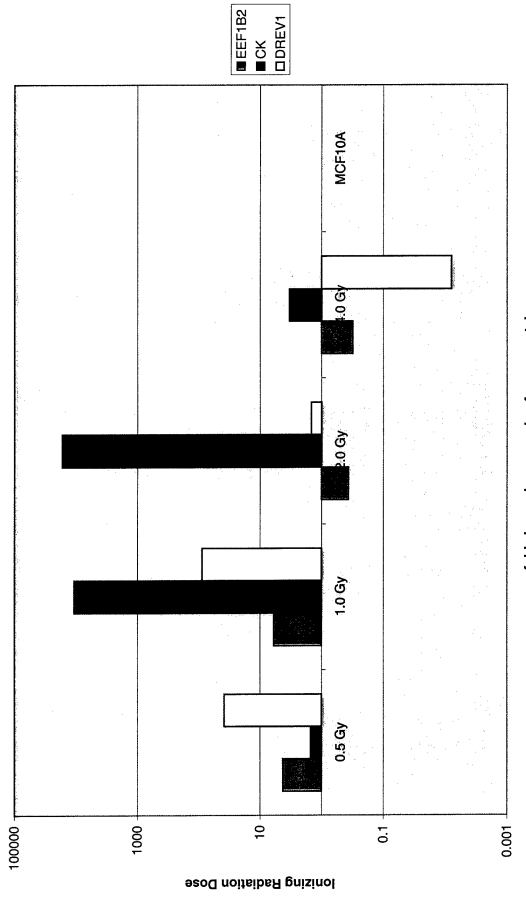


Relative mRNA expression levels



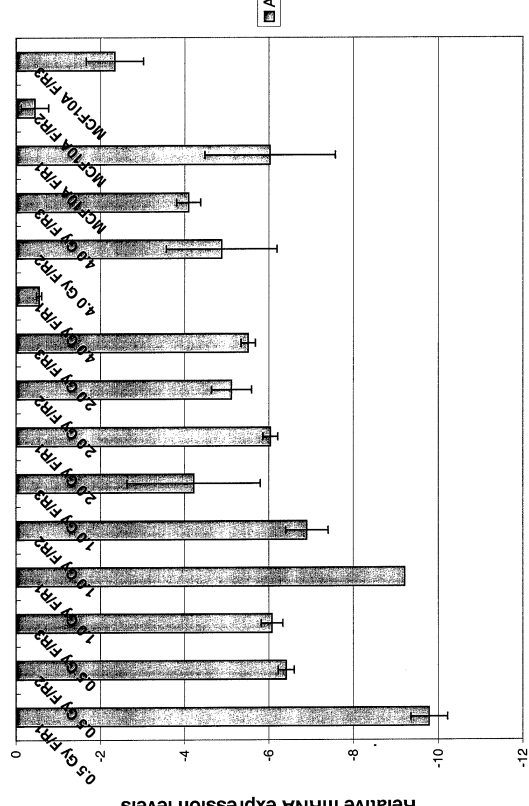
Ionizing Radiation Dose

N-fold difference from parental MCF10A



n-fold change in expression from parental

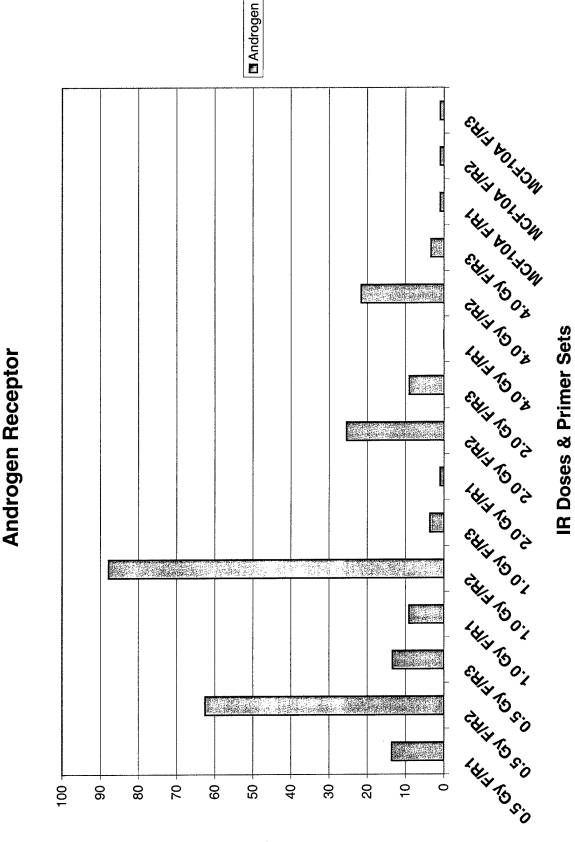
Relative mRNA expression levels



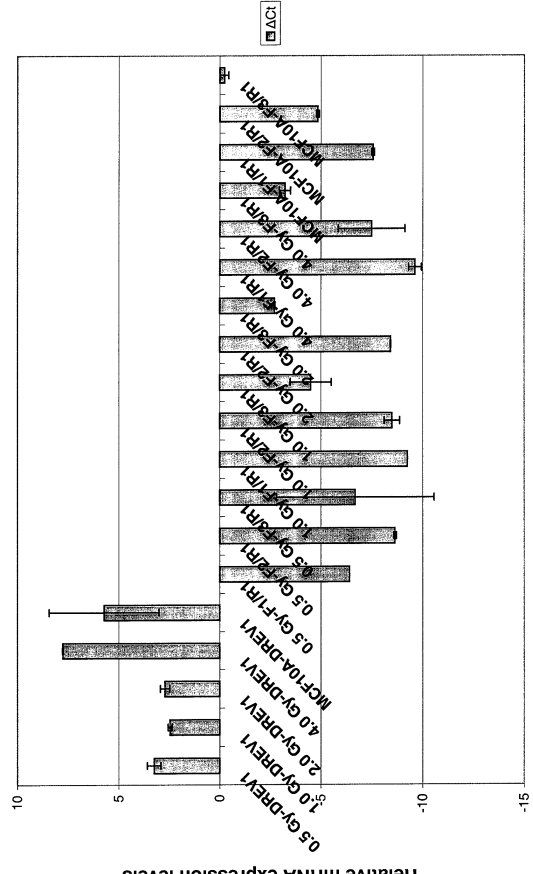
Androgen Receptor

🖪 Androgen

More in expression from parental MCF10A



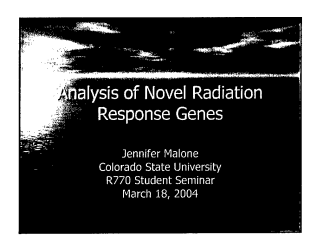


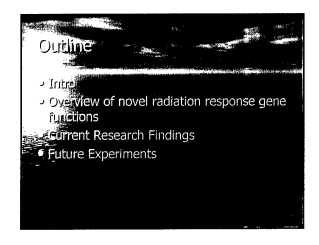


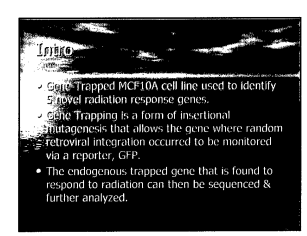
Mortal difference from parental MCF10A

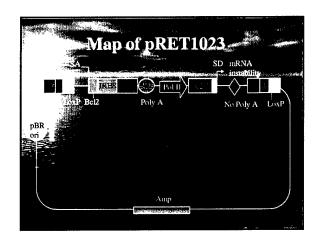


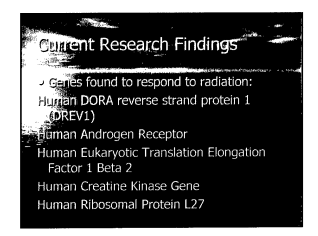
N-fold difference from MCF10A

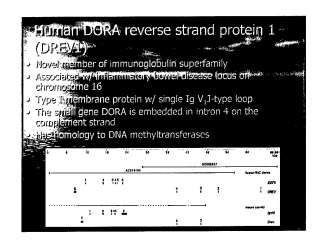


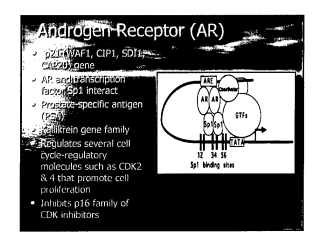


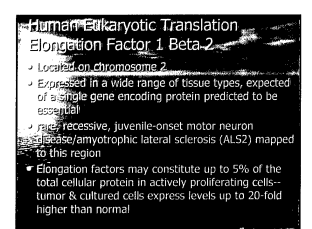


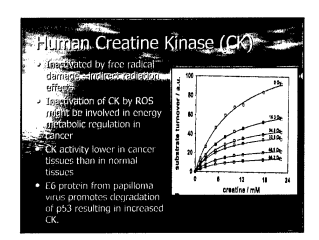


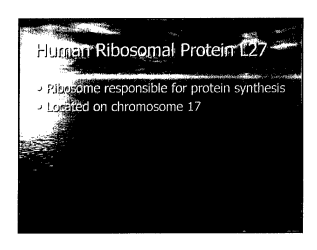


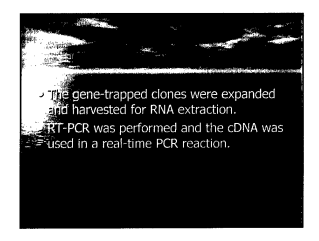


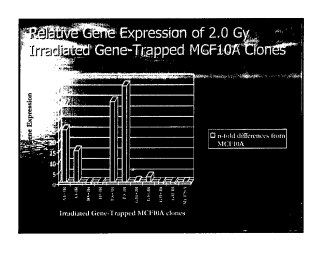


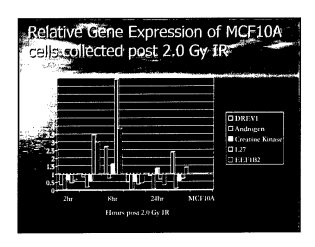


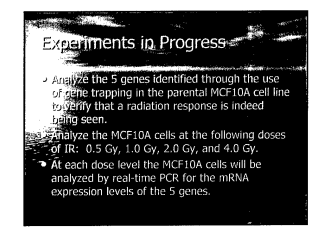


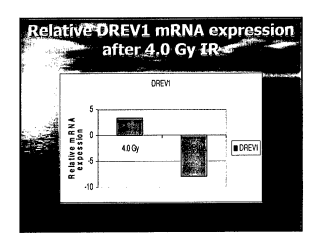


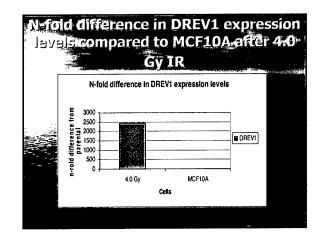


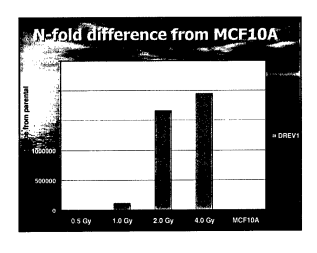


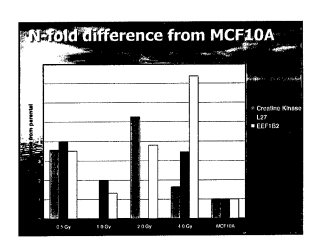


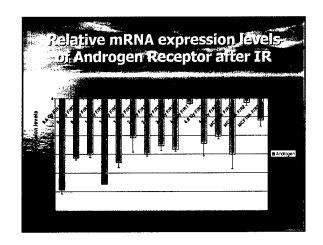


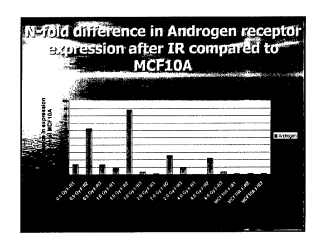


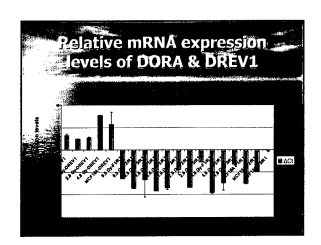


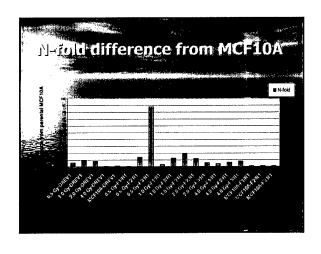






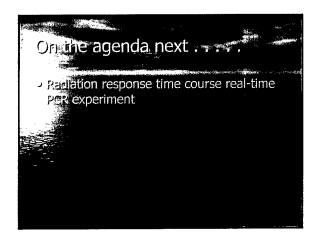






For each radiation response gene: at the IR dose with the highest response a time course of gene expression will be done at various time points following irradiation.

The time points will be at 2, 4, 8, 12, 24, and 30 hours post IR.



Acknowledgements

Pl. Robert Ullrich
Committee Members:
Dr. Sue Lana, Dr. Bill Hanneman, & Dr.
Mike Fox

Novel radiation response genes identified in MCF10A gene-trapped cells.

Jennifer Malone and Robert Ullrich

Department of Environmental and Radiological Health Sciences, Colorado State University Graduate Student, 491-7497, Jennifer.Malone@ColoState.EDU

Objective/Hypothesis: In this study, we have established an assay to identify novel genes that are affected by gamma irradiation and to characterize their function and role in early breast carcinogenesis. We hypothesize that the mutation of these genes or their abnormal expression in response to gamma irradiation is one of the causes of breast carcinogenesis.

Specific Aims: The specific aims of this study are:

1. To establish a high throughput assay for detection of variation in gene expression in human mammary epithelial cells using gene-trapped MCF10A clones; 2. To determine the effect of gamma irradiation on the expression of the reporter, green fluorescent protein (GFP); 3. To characterize the effect of gamma irradiation on the transformation of human mammary epithelial cells; 4. To identify the trapped genes affected by gamma irradiation in breast epithelial cells. **Methods:** We plan to establish an assay that will allow us to screen for breast cells that contain a single gene mutation using a technique called gene trapping. This will allow us to detect changes in the expression of a specific gene upon treatment with different doses of radiation. These radiation-responsive genes will be identified through the rapid amplification of cDNA ends (RACE) procedure and sequenced. Gene-trapped clones that are affected by radiation will be isolated and further analyzed by real-time PCR and compared to the parental to verify that a radiation response is being seen.

Results: The MCF10A gene-trapped library has been established and basal GFP levels have been measured. Gamma irradiation of the single cell gene-trapped clones at both 1.0 and 2.0 Gy has been performed. Clones that were either up- or down-regulated at least 2-fold in response to the radiation treatment have been expanded and analyzed by 3' RACE and sequencing. The five radiation response genes identified have been analyzed by real time PCR and cell cycle analysis. Study Design: Using the poly-A trap retrovirus RET, we have established a gene-trapped library of clones from human mammary epithelial cells (MCF10A). This provides a strong base for the identification of novel genes that may be involved in essential signaling pathways in human mammary epithelial cells. The reporter gene GFP, which has been incorporated into the genome of the cells, monitors the expression level of the endogenous trapped genes. We will compare basal GFP expression before and after exposure to varying low doses of gamma radiation (0-4 Gy) using replica plates of MCF10A gene-trapped clones. We will then identify the genes involved by using 3' RACE and sequencing. The identified radiation response gene's mRNA levels will be analyzed by real-time PCR analysis and compared to the parental MCF10A cell line after varying doses and time points following ionizing radiation.

Conclusions: This assay may prove to be a powerful tool in the identification of novel genes that are affected by gamma irradiation in the early stages of breast cancer progression. This study will provide new information on the effects of radiation-responsive genes that can lead to breast cancer as well as identify new markers for early detection of breast cancer. This study will focus on the identification of novel genes that are potential targets of gamma irradiation. It will provide essential information on the immediate and long-term effects of gamma irradiation of breast cells that may be the key to further understanding of the mechanism of radiation-induced breast cancer.



NOVEL RADIATION RESPONSE GENES IDENTIFIED IN MCF10A CELLS

3

Department of Radiological and Environmental Health Sciences Jennifer L. Malone and Robert L. Ullrich



Breast cancer may be induced with relatively high frequency by radiation. Ionizing radiation is one of the main returnent modellities used in the returnent of breast cancer. Induction is used in the returnent of breast cancer, leadering, and Hodglein's lymphoran to bill cancerous cells. While the use of medical radiation has undexchedly produced and the lives of many it is new wholens side offices. A radiation doscribated instruction that the control of the produced of the side of the sent in women. When a woman receives significant radiation prior to the age of 20, she becomes more likely to develop breast cancer. Thus there is considerable interest in understanding the cellular response to DNA champing spatis, performed by the considerable interest in understanding the cellular response to DNA champing spatis, perfordingly because the ability to deliver a curaint dose of radiation is frequently limited by the adverse meation of mortant issues within the radiation naturation and delong the companion of a cure, producing the publicage can be manipulated to improve the therapeutic ratio and hence, the chance of a cure.

We propose trait use expression to several unaxwore genes to measure agreement absorber that the depression of base genes may be one of the early steps in breast enteningements included by redialism. We have established an assay that allows us to severa for breast cells that centain a single gene mustican using a technique called gene trapping. Gene trapping is form of insertional managenesis specifically designed outlemying earl function by producing imagenic integration events. By employing the polyaderybation (poly A) In man park that unsersible from a selectable mearing enter deficing a poly A signal in a gener captures a poly A signal. Since poly the imagine in the sequence of the single security vector is stabilized only when a gene-imp vector captures a poly A signal. Since poly A intentified a signal expense, any gene could postentially be found that the sequence of the target genes, any gene could postentially be found that in a gene-imp regarder of the target genes, any gene could postentially be found in a distribution of could postentially be dependently of the expression of the target genes, any gene could postentially be found in a form that in a form that the substance of its transcripts in target cells. Open measurement with individual galaction, the adultion-response genes identified will be sequenced through the rapid amplification of cDNA ends (RACE) procedure. propose that the expression of several unknown genes is directly affected by gamma radiation.

<u>Hypothesis and Rationale</u> Mutainn of mood genes or their ahmanud expression in response to a single dose of gamma rodation is one of the causes of ourly broast carcinagenesis.

In experimental models, ionizing radiation incluses mammary transformation both in vivo and in vitro, beweet, the cellular and miscelular mechanisms of malation-include carcinogenesis are not known. To understand the mechanisms, it is necessary to determine the conditions that modular the susceptibility of this target tissue to foreign end misc. The assessment of the transforming allity containing radiation of breast application requires an abigity profilerating cell propalisien, which in turn has to cargies storate the east phenotypes. Since ionizing radiation induces features of neoplastic transformation in human breast cells, the identification of militaran phenotypes involved in breast cancer are of critical importance in understanding the publicancies of the disease.

The sim of our assay is to identify genes responsive to gamma imidation through the use of gene trapping. Gene trapping will lag the redustion-responsive genes and we will be able to monitor their expression levels using GPP. At the same time, we will be introducing a single allele gene disruption in the frame fairner-responsive gene by the imitogation of the problem of the discusse expression. We believe that by utilizing the gene carpoing exchange some of the unknown genes that are contributing some of milital and more familial levers an angel to the contributing some of the subney genes that are contributing some of the subney we genes that are contributing some of milital and more familial levers an argued to more also become exposed leading to manniers. It is of great importance that these radiation-induced mutuations be identified so that a secondary manner does not distribute our flower than these radiation-induced mutuations to them serve as markers for screening and hopefully aid it really described.

Methods

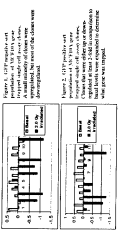
MCF-10A is a spontaneously non-tumorigenic immortalized luman breast epithelial cell line. We have generated RET infected GGH-Stream MCF-10A forces, which essentially are a gene-temporal liberay of mammary epithelia cells. This library can be used in identifying genese that are activated or inhibited in nammary epithelial cells in response to different geneous tegents or developmental signals. Since the infection net is one wire per cell lib library represents cells in which one functional gene is disrupted by the integration of the vector. It provides the strong base for the identification of novel genes that may be involved in essential signaling pathways in human mammary epithelial cells.

We have established a detection assay using the reporter, Green Fluorescent Protein (GFP), that has been incorparated into the genome of the clothes and will compare best singilated by the enrolegonus promoters of the trapped genes. We will compare basel GFP expression before and after exposure to warping tow does genome natistine (OF GS) using paging pulses of MCFI (DA clothes. We will then identify the genes involved by using a polymerase chain restion protocol (3 RACF) and esquencing malysis. Next, we will further characterize the clothes that are affected by genum airmitation by performing end time PCR to analyze gene expression, and again says to analyze anobrange-independent growth and tumorigenicity assays to confirm if the gene trap clotnes cause transformation.



Results

The gene-tapped MCT-10A clones were sorted by flow cytomerry into GFPs and
GFP- expression levels. The GFPs sort pool was then further sorted into GFP ight
and GFPs sorts. The GFP expression levels. Replica places were made for both the GFPs,
and GFPs sorts. The GFP expression level was measured tosting an ELISA assay both
before and after a 2.0 Gy close of FR.



3 RACE analysis was implemented to determine the identity of the trapped genes in the MCF10A ctones that were up- or down-regulated in response to a 2.0 Gy/dase of ion/zing radiation. The five genes identified through a BLAST homology search were.

Hunan Androgen Receptor . G.11.

Hunan Edaryoof Translation Elonguiton Factor 1 Beta 2: E8+
Hunan Creation Kinase Gene: B5+
Hunan Rhoscornal Protein L27, A4-Human DORA reverse strand protein 1 (DREV1); G10+

Quantitative real-time PCR analysis of nRNA expression of gone-trapped MCFIO14 doese installated with 2.000 yionizing patiests and MCFIO10. MCFIO10. The following epithelial cells harvested as various time points after 2.0 Gy doses. The following actualistic was done to obtain the results presented in the graphs below. 2-3-40.

This formula endoathers the relative gene expression by using the Ct values obtained from a PCR law time substanced print) excluded the property of the control of the point of the purple of the point of the purple of the endogenous control GAPDF to obtain the AACI values. MCFIOA permit cells as and callbrainer and set to 0.

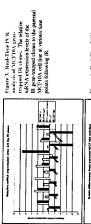


Figure 4. Real-Time FCR analysis of VIETER years-Trapped IR chours. The fold drange in mRNA expression level of the IR gene-trapped clones compared to the parental line. MCF10A.

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Cell cycle analysis was performed by flow cytometry to investigate if the DNA content varied duting planes of the CLP cycle after 2.0 Oyd ose of feating artifation. Analysis was performed on the purenal MCF10A cell fire without R and at varying time points after 2.2 Oyd dose of Rr. This arablysis was also performed on the five gene trapped clones both with and without 2.0 Oyd dose of RR. This acreptiones are done to disregard any flactuations in the mRNA expression levels being the lot a cell cycle phenometran.

Colorado Erres America Pares



Figure 6. Cell cycle analysis of DNA content in ploration crapture generatoryon cleans. The DNA content in Ol. O2, and Splase was performed by flow cytometry analysis. The generatopied clones were analysed with and without a 2.0 Gy dose of R.

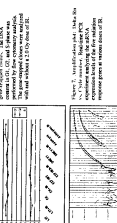


Figure 8. Real-Time IV IR analysis of DREVI mRN v expression. The N-fold difference in mRN acquession levels of DREVI in MCF10A cells with either a dose of 4.0 Gy or without IR.

N-fold difference from MCF10A

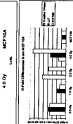


Figure 9. Real-Time IV R analysis of mRAN requession levels. The Nidol difference in mRN a repression levels of MCF10A at warons does of IR othe following genes: creatine kinase, phosomal profein L2, and ternelation choqual profein L2, and ternelation choqualon factor. Ivez 2.

Conclusions

This study identified five genes that are potential targets of gamma irradiation. The immediate and long-term effects of gamma irradiation to absence are little to be the key to further understanding the nerobanism of radiation-induced beast cancer. This seasy will also be useful for testing thet preducial environmental risk factors involved in breast carcinggenesis that may prove to be useful as markers for early detection of breast carcing and progress to be useful as markers for early detection of preast cancer and targets for thempeutic intervention.



This work was supported by a grant from the Department of Defense Breast Cancer Research Program, DAMD17-02-1-0349 to J. Malone





Novel radiation response genes identified in MCF10A gene-trapped cells.

Jennifer Malone and Robert Ullrich

Department of Environmental and Radiological Health Sciences, Colorado State University Abstract Category: Mutagenesis/Clastogenesis/Carcinogenesis

Objective/Hypothesis: In this study, we plan to establish an assay to identify novel genes that are affected by gamma irradiation and to characterize their function and role in early breast carcinogenesis. We hypothesize that the mutation of these genes or their abnormal expression in response to gamma irradiation is one of the causes of breast carcinogenesis.

Specific Aims: The specific aims of this study are:

1. To establish a high throughput assay for detection of variation in gene expression in human mammary epithelial cells using gene-trapped MCF10A clones; 2. To determine the effect of gamma irradiation on the expression of the reporter, green fluorescent protein (GFP); 3. To characterize the effect of gamma irradiation on the transformation of human mammary epithelial cells; 4. To identify the trapped genes affected by gamma irradiation in breast epithelial cells. **Methods:** We plan to establish an assay that will allow us to screen for breast cells that contain a single gene mutation using a technique called gene trapping. We will be able to detect changes in the expression of a specific gene upon treatment with different doses of radiation. These radiation-responsive genes will be identified through the rapid amplification of cDNA ends (RACE) procedure and sequenced. Gene-trapped clones that are affected by radiation will be isolated and further analyzed to see if the varying radiation doses can lead to malignant transformation.

Results: The MCF10A gene-trapped library has been established. Basal GFP levels have been measured. Gamma irradiation of the single cell clones at both 1.0 and 2.0 Gy has been performed. Clones that were either up- or down-regulated at least 2-fold in response to the radiation treatment have been expanded for RACE and sequencing analysis. The genes identified through sequencing have been analyzed by real time PCR.

Study Design: Using the poly-A trap retrovirus RET, we have established a gene-trapped library of clones from human mammary epithelial cells (MCF10A). It provides the strong base for the identification of novel genes that may be involved in essential signaling pathways in human mammary epithelial cells. We propose to establish a detection assay using the reporter gene GFP that has been incorporated into the genome of the cells, whose expression is regulated by endogenous promoters of the trapped genes. We will compare basal GFP expression before and after exposure to varying low doses of gamma radiation (0-2 Gy) using replica plates of MCF10A gene-trapped clones. We will then identify the gene(s) involved by using a polymerase chain reaction protocol and sequencing analysis. Next, we will further characterize the clones that are affected by gamma irradiation by performing colony formation assays (to determine survival), anchorage-independent growth and tumorigenicity assays on transformed clones that grow in soft agar.

Conclusions: This assay may prove to be a powerful tool in the identification of novel genes that are affected by gamma irradiation in the early stages of breast cancer progression. This study will provide new information on the effects of radiation-responsive genes that can lead to breast cancer as well as identify new markers for early detection of breast cancer. This study will focus on the identification of novel genes that are potential targets of gamma irradiation. It will provide essential information on the immediate and long-term effects of gamma irradiation of breast cells that may be the key to further understanding of the mechanism of radiation-induced breast cancer.

AACR Special Conference: Advances in Breast Cancer Research Abstract

Breast cancer may be induced with relatively high frequency by radiation. Ionizing radiation is one of the main treatment modalities used in the management of cancer. A radiation dose-related increase in the incidence of breast cancer has been seen in women. When a woman receives significant radiation prior to the age of 20, she becomes more likely to develop breast cancer. Thus there is considerable interest in understanding the cellular response to DNA-damaging agents, particularly because the ability to deliver a curative dose of radiation is frequently limited by the adverse reaction of normal tissues within the radiation treatment field. One approach to this problem is to understand the molecular mechanisms underlying the radiation responses of normal tissue so that critical molecular pathways can be manipulated to improve the therapeutic ratio and hence, the chance of a cure. We propose that the expression of several genes is directly affected by gamma radiation. Abnormal expression of these genes may be one of the early steps in breast carcinogenesis induced by radiation. We plan to screen breast cells that contain a single gene mutation using a technique called gene trapping. We will be able to detect changes in the expression of a specific gene upon treatment with different doses of radiation. These radiation response genes will be identified through the rapid amplification of cDNA ends (RACE) procedure and sequenced. Cells that are affected by radiation will be isolated and further analyzed to see if the changes can lead to the malignant transformation of the normal breast epithelial cell into a neoplastic cell. This assay may prove to be a powerful tool in the identification of novel genes that are affected by gamma irradiation in the early stages of breast cancer progression. This study will provide new information on the effects of radiation and genes that can cause breast cancer that are induced by radiation as well as identify markers for early detection of breast cancer and targets for therapeutic intervention.



IDENTIFICATION OF NOVEL GENES AFFECTED BY GAMMA IRRADIATION USING A GENE-TRAPPED LIBRARY OF HUMAN MAMMARY EPITHELIAL CELLS

Department of Radiological and Environmental Health Sciences Jennifer L. Malone and Robert L. Ullrich Colorado State University



Introduction

Breast cancer is one of the most common cancers among women and is the second leading cause of autoex death in women in the United States, executed only by Julia cancer. There are both genetic and environmental components associated with breast cancer. Genetic risks factors include mutations in supportants associated with breast cancer. Genetic risks factors include mutations in supportants is a known in risk and that the cancer. Genetic risks actions in the area radiation transmits, it is always in risk factor that can cause because cancer. Radiation therapy, used as part of breasts cancer, ring therapy for early breast cancer, is directed to normal breast tissue in order to endicate remaining malignant cells by inducing DNA damage and cell death.

We propose that the expression of several unknown genes is directly affected by gamma radiation. Abformul expression of these genes may be not of the early steps in breast carcinogenesis induced by radiation. We plan to exhibitish an assy than will allow us to serven for breast calls that cortain a single gene mutation sturgs a celahigate and lead to the server of the state of the depart of the expression of a specific gene upon teatment with different does of radiation. These radiation-responsive genes will be identified through the rapid amplification of chNA earls (RACE) procedure and sequenced. Cells that are affected by radiation will be subsidiated with the standard and further analyzed to see if the changes are lead to the multiparant transformation of the normal breast grithelia deal into a neoplastic cell. This steasy may prove to be a powerful tool in the identification of novel genes that are affected by gamma radiation and not early stages of breast cannor respectsion. This study will provide new information on the effects of multation and genes that can cause breast chance that are induced by spatial as identify markers and also also that cannor propersion. or early detection of breast cancer and targets for therapeutic intervention. Hypothesis and Rationale

Matation of novel genes or their abnormal expression in response to a single dose of gamma radiation is one of the causes of early breast carcinogenesis.

In experimental models, ionizing radiation incluses mammary transformation both in vivo and in vitro, betweeve, the collishest and moderation mechanisms of mediation-included carcinogeneis are not known. To understand the mechanisms, it is necessary to determine the conditions than modulate the susceptibility of this target tissue to originizing radiation. The assessment of the transforming ability of civinging radiation to be best applicate, which in turn has to express normal breast phenotypes. Since ionizing radiation incluses (loquers of neoplastic transformation in human breast earls, the identification of multipant phenotypes involved in breast cancer are of critical importance in adaptationing the publications of the disease.

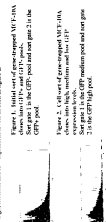
The aim of our assay is to identify genes responsive to gamma imadiation through the use of gene trapping. Gene trapping will ligh the radiation-responsive genes and we will be below to monitor their expression beeds using GPP. At the same time, we will be introducing a single alled gene disruption into the radiation-responsive gene by the integration of topoly At rap worker which will inderess expression. We believe that by utilizing the gene integration of the poly At rap worker without little areas expression. We believe that by utilizing the gene integrate schedule of the utilizing the state cancer can be identified. During radiation meatment of breast cancer some of the cells surrounding a urgested turns also become exponed leading to numbrains. It is of great impression the these radiation-induced mutations be identified so that a secondary cancer does not develop. The addiation-responsive genes identified can then serve as markers for screening and hopefully aid in early described.

Methods

MCF-10A is a spontaneously non-tumorigenic immortalized human breast epithelial cell line. We have alteredy generated RET-infected Calls-resistant MCF-10A clones, which essentially are a gen-tarapped library of mammary optibelial cells. At this point, this library can be used in identifying genes that are activated or inhibited in mammary epithelial cells in response to different genoxoxic agents or developmental signals. Since the infection mas is one vinue per cell, this library response cells in which one functional gare its disrepted by the integration of the vector. It provides the strong base for the identification of novel genes that may be involved in essential signaling pathways in human mammary epithelial cells.

We have established a detection assay using the reporter gene GFP (green fluorescent protein) that has been incorporated into the generon of the cells, whose expression is regulated by relogenous promosers of the turpung genes. We will compare based GFP expression before and after exposure to varying low dece general ending the control of the frapped genes. We will compare the stap grelles ablest to MFCFI to dates. We will estimate the control of genes involved the vising at polyments chain reaction protocol (3° RACE) and sequencing analysis. Next, we will further characterize the closes that are affected by garmai tradition by performing real-time PCR to analyze gene expression, so the garn reasys to canaly can undergoe independent growth and tumorigenicity assays to confirm if the gene trap closes cause transformation.

Results The great stayed MCF-10A closes were sorted by flow cytometry into GFP+ and GFPexpression it evelt, as soon in figure 1. The GFP+ sort pool was then further sorted into GFP high, medium and low expression levels as soon in figure 2.



Gene-trapped MCF10A-1023 cells were plated in 96-well plates to be replica plated in order to obtain single cell clones, not pooled clones. GFP expression was determined by a studivich ELISA protocol anneasured with a microplate reader. The GFP expression was visualized under a fluorescence microscope to verify the expression levels from the endogenous promoter of the trapped gene as either high medium one. As you can see below in figure 3, high, medium and low GFP expression was observed.



A. High GFP expressing M1023 gene-tapped clone from replica plate.

B. Medium GFP expressing M1023 gene-trapped clone from replica plate.

(1. Low GFP expressing M1023 clone from replica plate. ن

Analysis of GFP expression data from 2.0 Gy irradiated single cell MCF10A gene trapped clones by microplate reader.

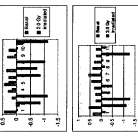


Figure 4. GFP negative sort population of MCF10A gene trapped single cell nessy clones. Sone clones were upregulated, but most of the clones were downregulated. Clones that were downregulated. Clones that were

either up or down-regulated at least 2-fold in comparison to basal levels were expanded to determine what gene was trapped.

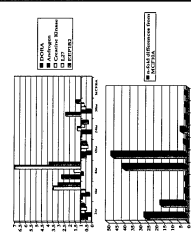
Figure 5. GFP positive sort population of MCF10A gene trapped single cell assay clones.

3' RACE analysis was implemented to determine the identity of the trapped genes in the MCFI0A clones that were up-or down-regulated in response to a 2.0 Gy doze of ionizing radiation. The six genes identified through a BLAST horselogy search were.

Human DORA reverse strand protein 1 (DREVI): G104
Human Adrogen Ceceptor. (211Human Edwarporic Translation Elongation Factor I Beta 2: E84Human Creation Kinase Gene: 184Human Nebound Protein L27: A4Human DNA sequence from close RP1290P20 on Chromosome 20: 114
Human DNA sequence from close RP1290P20 on Chromosome 20: 114

Quantitative real-time PCR analysis of mRNA expression of gene-trapped MCFIOA, elemest irradiated with 2.0 Gy storing relatives and MCFIOA element and MCFIOA elementalial cells harvested at various time points after 2.0 Gy does. The following calculation was done to obtain the results presented in the graphs below:

This formula calculates the relative gene expression by using the Ct values obtained from a PCR base line subtracted graph calculated by the iCycler software. The Ct values from the MCFIOA gene tapped clones were compared to the endogenous centred CAPUH to obtain the AACI values. MCFIOA purential cells were used as a calibrator and set to 1.



This study identified six genes that are potential targets of gamma irradiation. The irruncdiate and long-term effects of gamma irradiation on breast cells may be the key to further understanding the mechanism of radiation-induced breast cancer.
Establishment of his assay will also be useful for testing other potential
end of the common and the factors involved in breast carrioragenesis but may prove to be
useful as markers for early detection of breast cancer and targets for therapeanic Conclusions



This work was supported by a grant from the Department of Defense Breast Cancer Research Program, DAMD17-02-1-0349 to J. Malone



Real Time PCR analysis of gene-trapped MCF10A clones

Jennifer Malone Colorado State University October 23, 2003

Research Overview

- Hypothesis: Mutation of novel genes or their almormal expression in response to a single dose of gamma radiation is one of the causes of early breast carringenesis.
- Specific aim.1. To establish a high throughput assay for detection of variation in gene expression in human mammary epithelial cells using gene-trapped MCF-10A clones.
- Specific aim 2. To determine the effect of gamma irradiation on expression of reporter protein GFP.
- Specific aim_3. To identify the "trapped" genes affected by gamma irradiation.
- Specific aim 4. To characterize the effect of gamma irradiation on transformation of human mammary epithelial cells.

Breast Cancer & Radiation

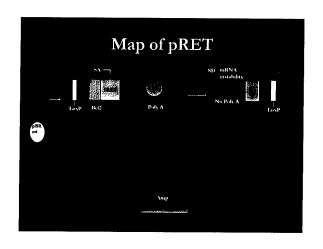
- Breast cancer may be induced with relatively high frequency by radiation.
- Ionizing radiation is one of the main treatment modalities used in the management of cancer.
- Radiation is used in the treatment of breast cancer, leukemia, and Hodgkin's lymphoma to kill cancerous cells.
- A radiation dose related increase in the incidence of breast cancer has been seen in women.
- When a woman receives significant radiation prior to the age of 20, she becomes more likely to develop breast cancer.

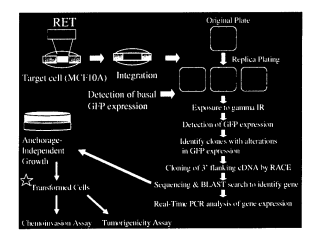
Gene Trapping

- Form of insertional mutagenesis.
- Disrupt gene function by intragenic integration.
- mRNA transcribed from a selectable marker gene lacking a poly A signal in a gene-trap vector is stabilized only when the gene-trap vector captures a cellular poly A signal.
- Poly A trapping occurs independently of the expression of target genes, regardless of its expression.
- The sequence of the 'trapped' gene can be identified using techniques that are based on the polymerase chain reaction (PCR), and this can lead to the isolation of novel genes regardless of their level of expression in vivo.

Materials

- MCF10A: immortalized human mammary epithelial cells that arose spontaneously.
- pRET: retroviral vector used for gene trapping. Contains a very strong splice acceptor and a poly A signal used to disrupt the trapped gene. Neo marker to select clones with integration and GFP for monitoring of endogenous trapped gene's expression level.





Sequencing Results

- 31 irradiated gene trapped clones sequenced
- Sequencing results plugged into BLAST
- 6 clones were homologous to known gene sequences:
-] : Human DORA reverse strand protein 1 (DREVI) : **G10+** Human Androgen Receptor : **G11-**Human Lukaryotic Translation Elongation Factor I <u>Beta 2</u> :

Human Creatine Kmase Gene : B5+

Human DNA sequence from clone RP1290F20 on Chromosome 20 : ${\bf H4\pm}$

The Next Step

- The six clones that had yielded homologous genes through BLAST search were analyzed.
- The clones were grown up and the cells were harvested for RNA extraction.
- RT PCR was performed and the cDNA was used in a real time PCR reaction.

Reverse Transcription

- 2.0 Gy irradiated gene trapped clones that had yielded homologous BLAST results were RT-PCR to analyze gene expression of the selected genes of interest
- Conditions:

Incubation: 25°C for 10 minutes

Reverse Transcription: 48°C for 30 minutes

RT Inactivation: 95°C for 5 minutes

Real Time PCR

■ Conditions:

UNG Incubation: 50 C for 2 minutes

AmpliTaq Gold Activation: 95 C for 10 minutes

PCR: started out at 40 cycles and increased up to 55

Denature: 95 C for 15 seconds

Anneal Extend: 60 C for I minute

Primers & Probes

- Three forward & reverse primers were designed by Primer Express for each gene of interest
- LapMan probes were designed by Primer Express for each gene of interest.
- Primer design requirements:

The $T_{\rm m}$ should be 58 to $60^{\rm o}{\rm C}$

Keep G C content in 20 80% range

The five nucleotides at the $\mathcal Y$ end should have no more than two G/C bases

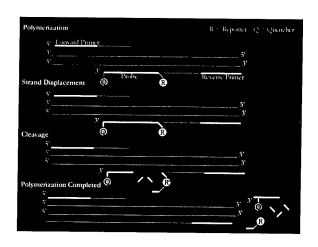
I orward & reverse primers should be as close as possible to the probe w_{\parallel} σ overlapping it

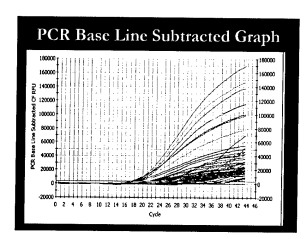
Avoid runs of in identical nucleoude especially G

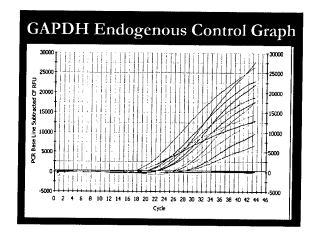
Probe design requirements:

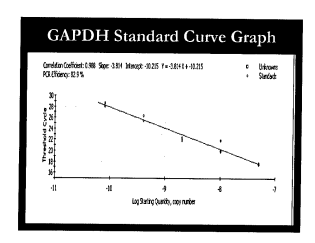
Avoid runs of an identical nucleotide, especially G. The 5' end of the probe cannot be a guanosine residu. The $V_{\rm m}$ should be 65 to 67°C.

TaqMan Probes The reporter dye FAM was used to label the 5' end of my gene specific probes The chromophore TAMRA was used to quench the probe on the 3' end

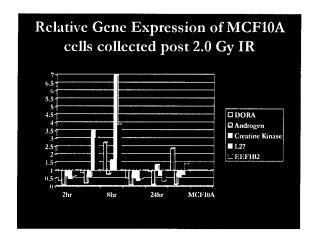


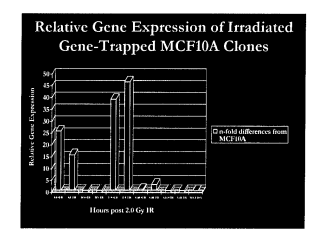


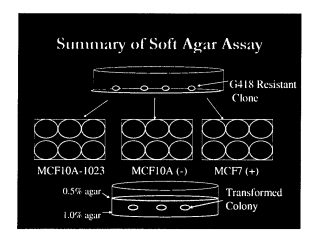




Data Calculations Absolute Standard Method for data quantification: ∆CT = CT (target) - CT (GAPDH) Comparative expression level for data quantification: = 2 △△CT







Future Directions

- Analyze the cell cycle distribution for my gene trapped irradiated clones and MCF10A various time points after IR by flow cytometry
- To determine if anchorage independent clones are fully malignant, MCF10A gene trapped clones will be injected subcutaneously into the subscapular area of 3 week old irradiated athymic female nude mice (BALB/c background).
- Characterize what other known proteins the radiation responsive gene(s) identified bind to

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